10/598,563 Page 3

5-7 7-8 8-9 9-10 10-11 11-12 12-13

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds :

1-2 1-6 2-3 3-4 4-5 5-6 11-12 12-13

exact bonds :

5-7 7-8 8-9 9-10 10-11

isolated ring systems :

containing 1 :

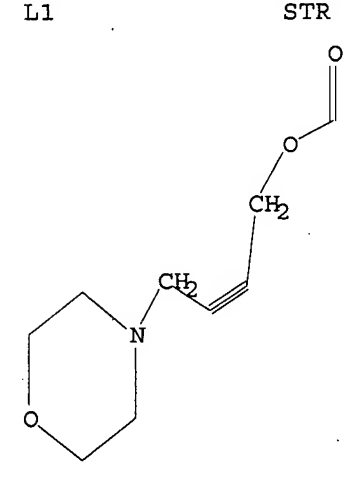
Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS 12:CLASS 13:CLASS

L1 STRUCTURE UPLOADED

=> d l1 L1 HAS NO ANSWERS



Structure attributes must be viewed using STN Express query preparation.

3 ANSWERS

=> s 11

SAMPLE SEARCH INITIATED 13:32:18 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 34 TO ITERATE

100.0% PROCESSED 34 ITERATIONS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 331 TO 1029

PROJECTED ANSWERS: 3 TO 163

L2 3 SEA SSS SAM L1

=> s l1 sss full

FULL SEARCH INITIATED 13:32:24 FILE 'REGISTRY'

Habte 10/29/2007

10/598,563 Page 4

FULL SCREEN SEARCH COMPLETED - 612 TO ITERATE

100.0% PROCESSED 612 ITERATIONS 64 ANSWERS

SEARCH TIME: 00.00.01

L3 64 SEA SSS FUL L1

=> file caplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 172.10 172.31

FULL ESTIMATED COST

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L4 38 L3

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10/598,563

L4 ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 2005:1103763 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 143:387062 Preparation of water soluble 4-amino-2-butynyl esters TITLE: having anticancer activity INVENTOR (S): Salama, Zoser B. PATENT ASSIGNEE(S): Germany

PCT Int. Appl., 92 pp. SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	PATENT NO.					KIND		DATE		APPLICATION NO.					DATE				
							_								-	-		-	
	WO	2005	5095369			A1		20051013		WO 2004-EP2090					20040302				
		W :	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
			CN.	co.	CR.	CU.	CZ.	DE,	DK,	DM,	DZ,	EC,	EE,	EG.	ES,	FI,	GB,	GD,	
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				A1 20061220				EP 2004-716240					20040302						
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										WO 2004-EP2090 W 20040302									

CASREACT 143:387062; MARPAT 143:387062 AB The present invention relates to water soluble 4-amino-2-butynyl or 4-(N-substituted amino)-2-butynyl esters (R1R2NCH2C.tplbond.CCH2O2R (I); variables defined below; e.g. 4-(morpholino)-2-butynyl acetate) and methods for production of said esters and the use of the esters for treatment

of cancer. 4-Morpholino-2-butynyl acetate and 4-morpholino-2-butynyl pivalate show the highest antitumor activity amongst 8 examples of I and low toxicity to fibroblasts. For I: R is H, a straight-chained or branched, (un)saturated aliphatic radical with 1-20 C-atoms ((un)substituted

≥1 times by C1-C6-alkyl, C1-C6-alkoxy, halogen, epoxy, amino, mercapto, a Ph ring ((un)substituted ≥1 times by C1-C6-alkyl, C1-C6-alkoxy, hydroxy, epoxy, amino, mercapto or halogen)), a cycloalkyl group with 4 to 7 atoms ((un)substituted ≥1 times by C1-C6-alkyl, C1-C6-alkoxy, hydroxy, epoxy, amino, mercapto or halogen); R1 and R2 are joined to form a heterocyclic ring with 3 to 6 C-atoms, (un) substituted ≥1 times by C1-C6-alkyl, C1-C6-alkoxy, hydroxy, halogen, epoxy, amino, mercapto, whereby at least one C-atom can be replaced by O, S or

or R1 and R2 = H, straight-chained or branched, (un)saturated aliphatic radical

with 1-20 C-atoms, (un)substituted ≥1 times by C1-C6-alkyl,

ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

106087-86-9 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, propanoate (ester) (9CI) (CA INDEX

866549-52-2 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, formate (ester) (9CI) (CA INDEX NAME)

866549-56-6 CAPLUS Cyclohexanecarboxylic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR

THIS

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) C1-C6-alkoxy, hydroxy, halogen, epoxy, amino, mercapto. Methods of prepn.

are claimed and .apprx.10 example prepns. are included. For example, 4-(morpholino)-2-butynyl acetate was prepd. in 2 steps starting from propargyl alc. and acetic acid to give propargyl acetate, which underwent a Mannich condensation with paraformaldehyde and morpholine in the presence of CuCl.

35956-47-9P, 4-Morpholino-2-butynyl acetate 35956-48-0P,

4- (Morpholino) -2-butynyl pivalate 54757-85-6P, 4- (Morpholino) -2-butynyl benzoate 106087-86-9P 4-(Morpholino)-2-butynyl propionate 866549-52-2P,

4-(Morpholino)-2-butynyl formate 866549-56-6P, 4-(Morpholino)-2-butynyl cyclohexanecarboxylate RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(drug candidate; preparation of water soluble 4-amino-2-butynyl esters having

anticancer activity)

35956-47-9 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX NAME)

35956-48-0 CAPLUS Propanoic acid, 2,2-dimethyl-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

54757-85-6 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, benzoate (ester) (9CI) (CA INDEX NAME)

L4 ANSWER 2 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:561937 CAPLUS

DOCUMENT NUMBER: 143:221810

TITLE: Virtual Screen for Ligands of Orphan G Protein-Coupled

Receptors

AUTHOR (S): Bock, Joel R.; Gough, David A. CORPORATE SOURCE

Department of Bioengineering, University of California

San Diego, La Jolla, CA, 92093-0412, USA

SOURCE: Journal of Chemical Information and Modeling (2005), 45(5), 1402-1414

CODEN: JCISD8; ISSN: 1549-9596

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB This paper describes a virtual screening methodol, that generates a

list of high-binding small mol. ligands for orphan G protein-coupled receptors (oGPCRs), circumventing the requirement for receptor three-dimensional structure determination Features representing the

based only on physicochem, properties of primary amino acid sequence, and ligand features use the two-dimensional atomic connection topol. and atomic

properties. An exptl. screen comprised nearly 2 million hypothetical oGPCR-ligand complexes, from which it was observed that the top 1.96% predicted affinity scores corresponded to "highly active" ligands against orphan receptors. Results representing predicted high-scoring novel ligands for many oGPCRs are presented here. Validation of the method was carried out in several ways: (1) A random permutation of the structure-activity relation of the training data was carried out; by comparing test statistic values of the randomized and non-shuffled data, we conclude that the value obtained with non-shuffled data is unlikely to have been encountered by chance. (2) Biol. activities linked to the compds. with high cross-target binding affinity were analyzed using computed log-odds from a structure-based program. This information was correlated with literature citations where GPCR-related pathways or processes were linked to the bioactivity in question. (3) Anecdotal, out-of-sample predictions for nicotinic targets and known ligands were performed, with good accuracy in the low-to-high "active" binding range. (4) An out-of-sample consistency check using the com. antipsychotic drug olanzapine produced "active" to "highly-active" predicted affinities for all oGPCRs in our study, an observation that is consistent with

documented findings of cross-target affinity of this compound for many different

GPCRs. It is suggested that this virtual screening approach may be used in support of the functional characterization of oGPCRs by identifying potential cognate ligands. Ultimately, this approach may have implications for pharmaceutical therapies to modulate the activity of faulty or disease-related cellular signaling pathways. In addition to application to cell surface receptors, this approach is a generalized strategy for discovery of small mols, that may bind intracellular enzymes and involve protein-protein interactions.

35956-47-9 RL: PAC (Pharmacological activity); PRP (Properties); BIOL (Biological

(virtual screen for ligands of orphan G protein-coupled receptors) 35956-47-9 CAPLUS

ANSWER 2 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 2-Butyn-1-ol, 4-(4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX NAME)

 $CH_2 - C = C - CH_2 - OAC$

REFERENCE COUNT:

108 THERE ARE 108 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 3 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) G-protein coupled receptor-ligand interactions and other biomol. interactions for drug design uses) 35956-47-9 CAPLUS

2-Butyn-1-ol, 4-(4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX NAME)

 $CH_2 - C = C - CH_2 - OAC$

L4 ANSWER 3 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:219996 CAPLUS

DOCUMENT NUMBER: 142:294328

Trainable system for predicting G-protein coupled TITLE: receptor-ligand interactions and other biomolecular

interactions for drug design uses Gough, David A.; Bock, Joel R. INVENTOR (S):

PATENT ASSIGNEE(S): USA U.S. Pat. Appl. Publ., 21 pp., Cont.-in-part of U.S. SOURCE:

Ser. No. 993,272. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

> PATENT NO. APPLICATION NO. KIND DATE DATE US 2005053999 Al 20050310 US 2004-973576 20041026 US 2002090631 US 2001-993272 A1 20020711 20011114 WO 2006057763 A2 20060601 WO 2005-US38693 20051025 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN. CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC. LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS. IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,

KG, KZ, MD, RU, TJ, TM PRIORITY APPLN. INFO.: US 2000-248258P P 20001114

> US 2001-993272 A2 20011114

> US 2004-973576 A 20041026

The invention is a teachable system and method for predicting the interactions of proteins with other proteins, nucleic acids and small mols. A database containing protein sequences and information regarding protein interactions is used to "teach" the machine. Proteins with unknown interactions are compared by the machine to proteins in the database. Homologs of proteins known to interact in the database are predicted to interact. The invention is used for anal. of protein-protein

interactions and protein-nucleic acid interactions, for prediction of protein epitopes, and for whole proteome interaction anal. Virtual

for ligands of orphan G-protein coupled receptors is provided. The method

of the invention can be used in drug design.

35956-47-9

RL: BSU (Biological study, unclassified); BIOL (Biological study) (predicted high-affinity ligands; trainable system for predicting

L4 ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1998:407803 CAPLUS DOCUMENT NUMBER:

129:81674

TITLE: Preparation and use of bi- and tricyclic pyridone derivatives against Alzheimer's disease

INVENTOR(S): Huber, Trottmann Gerda; Jakob-Roetne, Roland;

Kolczewski, Sabine: Norcross, Roger David; Woltering, Thomas Johannes

PATENT ASSIGNEE(S):

F. Hoffmann-La Roche A.-G., Switz. SOURCE: PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA?	CENT :	NO.					DATE			AP	PLICA	TION	NO.		ם	ATE	
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WO	9825	930			A2		1998	0618		WO	1997	-EP68	65		1	9971	209
WO	9825	930			A3		1998	0813									
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RIORITY	APP.	LN.	INPO	. :						EP	1996	-1200	50		4 1	9961	213
																•	
										ЕP	1997	-1156	14		4 1	9970	909
									1	US	1997	9765	41	1	43 1	9971	124

WO 1997-EP6865

W 19971209

OTHER SOURCE(S):

MARPAT 129:81674

L4 ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

The title compds. [1; A = H, C(0)R2, 3-cyclopropyl-1,2,4-oxadiazol-5-yl; R1 = (un) substituted Ph; R2 = lower alkyl, Q1-R5; Q1 = 0, NR6; R3, R4 =

R3R4 = SCH:CH; CH:CHS; CH:CHCH:CH, etc.; R5 = H, lower alkyl, lower alkenyl, etc.; R6 = H, lower alkyl, Ph, etc.], useful for the prophylaxis or treatment of illnesses which are connected with an inhibition of β -amyloid peptide activity, especially for the treatment of Alzheimer's disease, were prepared Thus, treatment of 8-(4-benzyloxyphenyl)-7-oxo-7H-

thieno[2,3-a]quinolizine-10-carboxylaic acid (preparation described)

with SOC12

in PhMe followed by reaction of the intermediate with 2-methoxyethylamine in dioxane afforded 74% the title compound II which showed IC50 of 22 µM against Aß production (measured using sandwich-ELISA in HEK cells).

209333-45-9P

RL: BAC (Biological activity or effector, except adverse); BSU

(Biological

study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation and use of bi- and tricyclic pyridone derivs. against

Alzheimer's disease) 209333-45-9 CAPLUS

7H-Thieno[2,3-a]quinolizine-10-carboxylic acid, 7-oxo-8-phenyl-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

ANSWER 5 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

L4 ANSWER 5 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:74807 CAPLUS

DOCUMENT NUMBER: 114:74807 TITLE:

Synthesis of acetylenic spirobutenolide derivatives and evaluation of their growth inhibitory effect on

cells in culture

Bador, P.; Chantepie, J.; Paris, J.; Quash, G. AUTHOR (5): CORPORATE SOURCE: Lab. Chim. Ther., Fac. Pharm., Lyon, F-69373, Pr. Arzneimittel-Porschung (1990), 40(10), 1135-9

SOURCE: CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE: Journal

LANGUAGE: English'

Acetylenic spirobutenolide amides and esters and their Mannich bases were synthesized to evaluate their growth inhibitory effect. The biol. tests used both normal and transformed cells and they show the selectivity of the prepared compds. The ester derivs, presented the best selectivity comparable to that of daunorubicin.

131967-24-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and growth inhibitory activity of, as acetylenic

spirobutenolide derivative)

131967-24-3 CAPLUS

1-Oxaspiro[4.5]dec-3-ene-4-carboxylic acid, 3-methyl-2-oxo-,

4-(4-morpholinyl)-2-butynyl ester, hydrochloride (9CI) (CA INDEX NAME)

● HCl

IT 131926-46-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antitumor acetylenic spirobutenolide derivs.)

131926-46-0 CAPLUS

1-Oxaspiro(4.5)dec-3-ene-4-carboxylic acid, 3-methyl-2-oxo-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 6 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:571476 CAPLUS

DOCUMENT NUMBER: 113:171476

TITLE: Preparation of butynylamine derivatives for treatment of pollakiuria and like diseases

INVENTOR (S): Kimura, Kiyoshi; Kise, Masahiro; Morita, Iwao

PATENT ASSIGNEE(S): Nippon Shinyaku Co., Ltd., Japan

Brit. UK Pat. Appl., 40 pp CODEN: BAXXDU

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT INFORMATION:					
PATENT NO.	KIND		APPLICATION NO.		DATE
GB 2222828	A	19900321	GB 1989-20766		19890913
GB 2222828	В	19920429			
IL 91377	A		IL 1989-91377		19890822
CN 1041582	A		CN 1989-106930		
CN 1038410	B	19980520			
EP 359311		19900321	EP 1989-202235		19890905
EP 359311		19910703			
EP 359311		19970115			
R: DE, IT, NL,					
ES 2016060	A6	19901001	ES 1989-3097		19890912
KR 154325			KR 1989-13217		
JP 02218651	A				
JP 06069996					,
	A2	19920128	HU 1989-4825		19890913
CH 680440	A5	19920831			
CA 1317943		19930518			
FR 2639044	Al	19900518	FR 1989-12032		
FR 2639044		19930806			17070314
US 5036098		19910730	US 1989-407228		19890914
BE 1003256		19920211			
US 5036098		19931102	US 1992-90002826		
PRIORITY APPLN. INFO.:		.,,,,,,,,	JP 1988-231272		19880914
			J. 1900-1311/2	^	. 7000914
			US 1989-407228	A	19890914

A 19890914

CASREACT 113:171476; MARPAT 113:171476 AB The title derivs. of the R1R2C(OH)COACR3R4C.tplbond.CCH2NR5R6 (R1, R2 = cycloalkyl, Ph, or 2-thienyl; R3, R4 = H, alkyl, or together with the adjacent C form a cycloalkyl; R5, R6 = H, alkyl; or together with the N form a cyclic amino; A = 0 or NR where R = H or alkyl), and their pharmacol. acceptable salts, are prepared. The derivs. show

and Ca2+ antagonism. Thus, to a heated mixture containing

1,1-dimethy1-2~

anticholinergic

propynyl α -cyclohexyl- α -phenylglycolate, paraformaldehyde, and CuCl in dioxane was added Et2N to give 4-diethylamino-1,1-dimethyl-2butynyl a-cyclohexyl-a-phenylglycolate (I), which was isolated as the HCl salt. I-HCl showed PA2 values for anticholinergic action and Ca2+ antagonism upon detrusor muscles of excised rabbit bladder of 7.33 and 6.72, resp. 129927-39-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of, as anticholinergic and calcium antagonist) 129927-39-5 CAPLUS

ANSWER 6 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 2-Thiopheneacetic acid, \alpha-hydroxy-\alpha-2-thienyl-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

ANSWER 7 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

L4 ANSWER 7 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1989:57493 CAPLUS DOCUMENT NUMBER: 110:57493 TITLE: Heterocyclylalkylphenyl N-phenylcarbamate derivatives as acetylcholinesterase inhibitors, their preparation, and formulations containing them INVENTOR (S): Tamura, Toshiya; Tsukamoto, Shinichi; Usuda, Shinji; Harada, Masatomi PATENT ASSIGNEE (S): Yamanouchi Pharmaceutical Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 40 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE

PATENT NO. KIND APPLICATION NO. DATE ------------JP 63170356 19880714 JP 1986-311852 19861230 A PRIORITY APPLN. INFO.: JP 1986-311852 19861230

OTHER SOURCE(S): MARPAT 110:57493

The title compds. I (R1 = H, halo, lower alkoxy, OH, etc.; R2 = H, lower alkyl; NR3R4 = (substituted) heterocyclyl which may be fused to a benzene ring; n = 1-5; Y = 0, S, imino; X = CH2CH:CH, CH2C.tplbond.C, etc.], useful as acetylcholinesterase inhibitors, were prepared. A mixture of m-(piperidinomethyl)phenol and m-ClC6H4NCO in C6H6 was refluxed for 2 h

give carbamate II. II in vitro exhibited an IC50 of 0.12 µM against acetylcholinesterase.

118511-48-1P IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as acetylcholinesterase inhibitor)

118511-48-1 CAPLUS

Carbamic acid, (3-chlorophenyl)-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 8 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN

ACCESSION NUMBER:

1988:406012 CAPLUS DOCUMENT NUMBER: 109:6012

TITLE: Carbon-13 NMR study of some acetylenic amines, their N-oxides and their rearrangement products

AUTHOR (S): Al-Rawi, Jasim M. A.; Khuthier, Abdul-Hussain;

Abachi,

CORPORATE SOURCE:

Coll. Sci., Univ. Mosul, Mosul, Iraq SOURCE: Spectrochimica Acta, Part A: Molecular and

Biomolecular Spectroscopy (1987), 43A(9), 1121-3

CODEN: SAMCAS; ISSN: 0584-8539 DOCUMENT TYPE: Journal

LANGUAGE:

English CASREACT 109:6012 OTHER SOURCE(S):

$$ACOCH_2C = C = CH_2$$

$$ACOCH_2C = CCH_2N$$

$$X$$

$$I$$

The 13C NMR spectra of acetylenic amines I (X = a bond, CH2, CHMe, O, CH2CH2, etc.) and some of their N-oxides were analyzed. Thermal

rearrangement of the oxides gave allenes (II). 35956-47-9P IT

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation, carbon-13 NMR and oxidation of) 35956-47-9 CAPLUS

CN 2-Butyn-1-ol, 4-(4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX NAME)

CH2-C=C-CH2-OAC

114906-22-8P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, carbon-13 NMR and rearrangement of)

114906-22-8 CAPLUS 2-Butyn-1-ol, 4-(4-oxido-4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX

ANSWER 8 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

ANSWER 9 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) CM 2

CRN 144-62-7

CMF C2 H2 O4

L4 ANSWER 9 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1987:515227 CAPLUS DOCUMENT NUMBER: 107:115227 TITLE: Acetylenic amines of potential pharmacological value AUTHOR (S): Abachi, F. T.; Yousif, W. H.; Al-Rawi, M. M.; Khodr, A. M.; Khuthier, A. H. CORPORATE SOURCE: Coll. Vet. Med., Univ. Mosul, Mosul, Iraq SOURCE: Journal of the Iraqi Chemical Society (1986), 11(1), 105-14 CODEN: JICSDK; ISSN: 0379-8321 DOCUMENT TYPE: Journal LANGUAGE: English OTHER SOURCE(S): CASREACT 107:115227 AB Amines R1CH2C.tplbond.CCH2NR2 [R1 = MeO, 3,5-(02N)2C6H3C02; NR2 = piperidino, morpholino, 4-formyl-1-piperazinyl], which showed mydriatic activity and its usefulness in the treatment of Parkinsonism, were prepared by the Mannich reaction. 110197-02-9P 110197-03-0P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation and pharmacol. activity of) 110197-02-9 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, 3,5-dinitrobenzoate (ester) (9CI) (CA

NO2 110197-03-0 CAPLUS 2-Butyn-1-ol, 4-(4-morpholinyl)-, 3,5-dinitrobenzoate (ester), ethanedicate (1:1) (salt) (9CI) (CA INDEX NAME) CM

CRN 110197-02-9 CMF C15 H15 N3 O7

L4 ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1987:32952 CAPLUS 106:32952 DOCUMENT NUMBER:

TITLE: Synthesis and properties of acetylenic amino esters of

some aliphatic acids AUTHOR (S):

Ergashev, M. S.; Kasymova, S. S.; Kulakhmatova, M. A. CORPORATE SOURCE: Tashk. Gos. Univ., Tashkent, USSR SOURCE: Izvestiya Vysshikh Uchebnykh Zavedenii, Khimiya i

Khimicheskaya Tekhnologiya (1986), 29(1), 39-41-CODEN: IVUKAR; ISSN: 0579-2991

DOCUMENT TYPE: Journal

LANGUAGE: Russian OTHER SOURCE(S): CASREACT 106:32952

GΙ

Morpholinobutynyl alkanoates I (R = C2-C9 n-alkyl) were prepared in 72.4-83.2% yields by Mannich reactions of morpholine and paraformaldehyde with RCO2CH2C.tplbond.CH in dioxane containing CuCl.

IT 106087-86-9P 106087-87-0P 106087-88-1P 106087-89-2P 106087-90-5P 106087-91-6P

106087-92-7P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 106087-86-9 CAPLUS

2-Butyn-1-ol, 4-(4-morpholinyl)-, propanoate (ester) (9CI) (CA INDEX

106087-87-0 CAPLUS Butanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 106087-88-1 CAPLUS

Pentanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

106087-89-2 CAPLUS Hexanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

106087-90-5 CAPLUS Octanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

106087-91-6 CAPLUS Nonanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 11 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1987:27650 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

106:27650

Synthesis and hypocholesterolemic activity of TITLE:

aminobutynyl linoleates Ergashev, M. S.; Makhsumov, A. G.; Khadzhiev, A. K. AUTHOR(S): CORPORATE SOURCE: Med. Inst., Tashkent, USSR Khimiko-Farmatsevticheskii Zhurnal (1986), 20(9),

SOURCE: 1050-1

CODEN: KHFZAN; ISSN: 0023-1134

DOCUMENT TYPE: Journal

LANGUAGE: Russian

AB Me(CH2)4CH:CHCH2CH:CH(CH2)7CO2CH2C.tplbond.CCH2NR2 (I, R = Et or CH2Ph or NR2 = piperidinyl or morpholinyl) were prepared from propargyl linoleate [106059-79-4], CH2O and the appropriate amine. In studies in rabbits with

exptl. atherosclerosis and hypercholesterolemia, I (NR2 = morpholino) [106059-82-9 and I (R = CH2Ph) [106059-83-0] were more active as hypocholesterolemics than were the other 2 compds. All were more

effective than the hypocholesterolemic Arakhides. 106059-82-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation and hypocholesterolemic activity of) 106059-82-9 CAPLUS

9,12-Octadecadienoic acid (92,122)-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

Double bond geometry as shown.

ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

106087-92-7 CAPLUS

Decanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 12 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1987:4504 CAPLUS

DOCUMENT NUMBER: 106:4504

TITLE: Amino ester acetylene derivatives of sorbic acid Makhsumov, A. G.; Tadzhibaev, U.; Ergashev, M. S. AUTHOR (S): CORPORATE SOURCE: Tashk. Gos. Med. Inst., Tashkent, USSR

Uzbekekii Khimicheskii Zhurnal (1985), (5), 63-5 SOURCE:

CODEN: UZKZAC; ISSN: 0042-1707 DOCUMENT TYPE: Journal

LANGUAGE: Russian

AB Mannich reaction of propargyl sorbate with R2NH (R = hexyl, octyl, PhCH2; R2N = morpholino, anabasino, cytisino) and paraform in dioxane containing

Cu(OAc)2 at 100-105° gave 6 corresponding Me(CH:CH)2CO2CH2C.tplbond.CCH2NR2 in 76.1-92.1% yield.

105566-28-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by Mannich reaction of propargyl sorbate)

105566-28-7 CAPLUS 2,4-Hexadienoic acid, 4-(4-morpholinyl)-2-butynyl ester, (E,E)- (9C1)

INDEX NAME)

Double bond geometry as shown.

L4 ANSWER 13 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1984:406587 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 101:6587

Hypolipidemic activity of derivatives of propargyl TITLE:

esters of linolenic acids AUTHOR (S):

Makhaumov, A. G.; Khadzhiev, A. K.; Gul'mirzaeva, I. K.; Khadzhiev, K. Kh.; Ergashev, M. S.; Madikhanov,

CORPORATE SOURCE:

SOURCE: Piziol. Aktiv. Veshchestva (1983), (15), 62-6

From: Ref. Zh., Khim. 1984, Abstr. No. 3Zh107 DOCUMENT TYPE: Journal

LANGUAGE: Russian

CASREACT 101:6587 OTHER SOURCE(S): AB Title only translated.

IT 90430-69-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and hypolipidemic activity of)

90430-69-6 CAPLUS

CN 9,12,15-Octadecatrienoic acid, 4-(4-morpholinyl)-2-butynyl ester, (Z, Z, Z) -

(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L4 ANSWER 15 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1975:409070 CAPLUS

DOCUMENT NUMBER: 83:9070

Synthesis of y-substituted propargyl alcohols,

TITLE: their ethers and esters

Kruglikova, R. I.; Berestevich, B. K.; Babaeva, L. AUTHOR (S):

G.;

Mosk. Inst. Tonkoi Khim. Tekhnol. im. Lomonosova, CORPORATE SOURCE:

Izvestiya Vysshikh Uchebnykh Zavedenii, Khimiya i SOURCE:

Khimicheskaya Tekhnologiya (1974), 17(12), 1824-7

CODEN: IVUKAR; ISSN: 0579-2991

DOCUMENT TYPE: LANGUAGE: Russian

AB RC.tplbond.CCH2OH (R = Me, MeOCH2, CH2:CH, Ph, Me2NCH2, Me2C(OH), 1-hydroxycyclohexyl, PhCH(OH)) were prepared in 38-59% yield. E.g., H2C:CH2C.tplbond.CCH2OH was prepared by treatment of HC.tplbond.CCH:CH2

with EtMgBr, followed by HCHO. R1C.tplbond.CCH2OMe [R1 = H, Me, MeOCH2, Ph, Me2NCH2, MeCO2CH2, C1CH2, BrCH2, MeC(OH)] were prepared in 39-85% yield, usually by methylation of the resp. alcs. RC.tplbond.CCH2O2CC6H4NO2-p (R ■ H, Me, MeOCH2, Ph, Me2NCH2, Br) and RC.tplbond.CCH2O2CPh (R = H, Me, MeOCH2, CH2:CH, Ph, 1-hydroxycyclohexyl, Me2NCH2, Et2NCH2,

piperidinomethyl, morpholinomethyl) were prepared by standard methods. 54757-85-6P 54757-94-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 54757-85-6 CAPLUS

RN 2-Butyn-1-ol, 4-(4-morpholinyl)-, benzoate (ester) (9CI) (CA INDEX NAME)

54757-94-7 CAPLUS Morpholinium, 4-[4-(benzoyloxy)-2-butynyl]-4-methyl-, iodide (9CI) (CA INDEX NAME)

Habte

L4 ANSWER 14 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1979:125487 CAPLUS 90:125487 DOCUMENT NUMBER:

TITLE: Study of the inhibiting properties of

1-chloro-2-oxo-3-oxa-5-hexyne and its amino derivatives in the acid corrosion of metals Tsalikova, Z. M.; Karaev, S. F.; Shikhiev, I. A.;

Asadullaev, A. F. CORPORATE SOURCE: Azerb. Inst. Nefti Khim., Baku, USSR

SOURCE: Azerbaidzhanskii Khimicheskii Zhurnal (1978), (3),

CODEN: AZKZAU; ISSN: 0005-2531

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB The inhibiting effects of RCH2CO2C.tplbond.CCH2R1 (I; R = H, Cl, Et2N, Bu2N, piperidino, or morpholino; R1 = H, Et2N, Bu2N, piperidino, or morpholino) on the corrosion of St. 3 [39296-41-8] in 4 N HCl at 60° were studied. In general, the corrosion inhibiting effect was decreased, compared to I (R = Cl, R1 = H) [627-09-8], with introduction of an amino substituent in the acetate moiety; i.e., I (R = Et2N, piperidino, or morpholino, R1 - H); however, the greatest inhibiting effect was exhibited by I (R = Bu2N, RI = H) (54480-21-6). Introduction of 2 amino substituents decreased the inhibiting effect by a factor of .apprx.2.

54928-26-6

AUTHOR (S):

RL: USES (Uses) (corrosion inhibition by, of steel in hydrochloric acid solution)

54928-26-6 CAPLUS

4-Morpholineacetic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 15 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

(Continued)

L4 ANSWER 16 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1975:149868 CAPLUS 82:149868 DOCUMENT NUMBER: TITLE: Physiological activity of new aminoacetylenic sorbic acid esters AUTHOR(S): Abdullaev, Sh. U.; Makhsumov, A. G.; Usmanov, M. Tashk. Gos. Univ., Tashkent, USSR CORPORATE SOURCE: Dokl. Vses. Konf. Khim. Atsetilena, 4th (1972), SOURCE: Meeting Date 1972, Volume 1, 500-3. Editor(s): Azerbaev, I. N. Akad. Nauk Kaz. SSR, Inst. Khim. Nauk: Alma-Ata, USSR. CODEN: 30AKA7 DOCUMENT TYPE: Conference LANGUAGE: Russian AB Iodomethylates of 6 sorbic acid aminoacetylenic esters showed bactericidal activity against 7 pyrogenic and intestinal bacterial species. Even the most active of these compds., sorbic acid 4-(N-3-methylpiperidinobut-2ynyl) ester iodomethylate [54951-08-5], was somewhat less effective than several commonly used antibiotics. 54951-10-9 RL: BAC (Biological activity or effector, except adverse); BSU study, unclassified); BIOL (Biological study) (bactericidal activity of) 54951-10-9 CAPLUS Morpholinium, 4-methyl-4-(4-((1-oxo-2,4-hexadienyl)oxyl-2-butynyl)-, iodide, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

• I -

L4 ANSWER 18 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1973:536947 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 79:136947 Sorbates of acetylenic amino alcohols TITLE: Makheumov, A. G.; Abdullaev, Sh. U. AUTHOR (S): CORPORATE SOURCE: SOURCE: Khim, Atsetilena Tekhnol, Karbida Kal'tsiya (1972) From: Ref. Zh., Khim. 1973, Abstr. No. 9Zh368 DOCUMENT TYPE: Journal LANGUAGE: Russian Paraformaldehyde (0.15 mole), 0.12 mole piperidine, 0.1 mole propargyl sorbate, CuCl, and dioxane was heated 7 hr at 94-6°, and the product converted into the methiodide to give 90% MeCH:CHCH:CHCO2CH2C.tplbond.CCH2R.MeI (I) (R = piperidino). Other I prepared were (R and & yield given): 2-methylpiperidino, 87.5; 3-methylpiperidino, 88.2; 4-methylpiperidino, 87; 5-ethyl-2methylpiperidino, 75.6; morpholino, 91.3; 2-(3-pyridyl)piperidino, 85; and hexahydroazepino, 85. ΙŢ 50669-11-9P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 50669-11-9 CAPLUS RN Morpholinium, 4-methyl-4-[4-[(1-oxo-2,4-hexadienyl)oxy]-2-butynyl]-, iodide (9CI) (CA INDEX NAME)

L4 ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1975:124676 CAPLUS DOCUMENT NUMBER: 82:124676 TITLE: Reaction of 2-propyn-1-ol chloroacetate with amines AUTHOR (S): Karaev, S. F.; Tsalikova, Z. M.; Shikhiev, I. A. CORPORATE SOURCE: Azerb: Inst. Nefti Khim, im. Azizbekova, Baku, USSR Azerbaidzhanskii Khimicheskii Zhurnal (1974), (4), SOURCE: CODEN: AZKZAU; ISSN: 0005-2531 DOCUMENT TYPE: Journal LANGUAGE: Russian The reaction of ClCH2COCl with HOCH2C.tplbond.CH gave HC.tplbond.CCH2O2CCH2C1, which was aminated with HNR2 to give HC.tplbond.CCH2O2CCH2NR2 (I, NR2 = NEt2, NBu2, piperidino, morpholino). II were aminomethylated with paraformaldehyde in HNR12 to give R12NCH2C.tplbond.CCH2O2CCH2NR2 (R12N, R2N given): Et2N, Et2N; bu2N, Bu2N; piperidino, piperidino; morpholino, morpholino; piperidino, Et2N. 54928-26-6P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 54928-26-6 CAPLUS 4-Morpholineacetic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA

L4 ANSWER 19 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1973:71357 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 78:71357 TITLE: Synthesis and properties of acetylenic amino esters of palmitic and stearic acids AUTHOR(S): Abdurakhimov, A.; Maksumov, A. G.; Il'khamdzhanov, P. CORPORATE SOURCE SOURCE: Tr. Inst. Khim. Nefti Prirod. Solei, Akad. Nauk Kaz. SSR (1971), No. 3, 145-9 From: Ref. Zh., Khim. 1972, Abstr. No. 42h164 DOCUMENT TYPE: Journal LANGUAGE: Russian AB Mannich reaction of propargyl esters of palmitic and stearic acids in dioxane with Cu212 [better than Cu(AcO)2, Cu2Cl2, CuCl2, Cu2Br2, or CuBr2] as catalyst gave Me(CH2)nCO2CH2C.tplbond.CCH2Z (Z and % yield for n = 14 and n = 16 given): morpholino, 79, 79.6; piperidino, 79.6, 82.8; 2-methylpiperidino, 66.3, 74.9; 3-methylpiperidino, 70, 77.9; 4-methylpiperidino, 78.8, 80.1; 5-ethyl-2-methylpiperidino, 66.1, 69.8; 2-(3-pyridyl)piperidino, 78.9, 79.4; Me2N, 74.4, 67.9; Et2N, 70.8, 82.2; Bu2N, 79.3, 82.1; and Bz2N (sic), 86.7, 87.4. 29237-96-5P 38022-01-4P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 29237-96-5 CAPLUS Octadecanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX

RN 38022-01-4 CAPLUS
CN Hexadecanoic acid, 4-(4-morpholinyl)-2-butynyl eater (9CI) (CA INDEX NAME)

L4 ANSWER 20 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1972:526576 CAPLUS DOCUMENT NUMBER: 77:126576

ORIGINAL REFERENCE NO.: 77:20853a,20856a

Condensation of propargyl palmitate with amines TITLE: AUTHOR (S): Abdurakhimov, A.; Makhaumov, A. G.; Safaev, A. S.;

Il'khamdzhanov, P. CORPORATE SOURCE: USSR

SOURCE: Tr. Tashkent. Politekh. Inst. (1970), No. 64, 29-32

Prom: Ref. Zh., Khim. 1971, Abstr. No. 22Zh230

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB The maximum yield is obtained in the title reaction if HCHO is used, rather

than (HCHO)x, and Cu(OAc)2 is used as catalyst. Thus, 0.015 mole 40% HCHO, 0.01 mole piperidine, 0.01 mole Me(CH2)14CO2CH2C.tplbond.CH, 40 ml dioxane, and 0.15 g Cu(OAc)2 was heated 6 hr at 96-8° to give 83% Me(CH2)14CO2CH2C.tplbond.CCH2R (R = piperidino). Similarly prepared was

82.8% morpholino analog. IT 38022-01-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 38022-01-4 CAPLUS

Hexadecanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX

L4 ANSWER 22 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1970:487375 CAPLUS

DOCUMENT NUMBER: 73:87375 ORIGINAL REFERENCE NO.: 73:14280h,14281a

Derivatives of stearic acid aminoester acetylenes TITLE:

Il'khamdzhanov, P.; Makhsumov, A. G.; Absurakhimov, AUTHOR (S):

SOURCE:

Zhurnal Prikladnoi Khimii (Sankt-Peterburg, Russian Pederation) (1970), 43(6), 1414-15

CODEN: ZPKHAB; ISSN: 0044-4618

DOCUMENT TYPE: Journal LANGUAGE: Russian

Me(CH2)16CO2CH2C.tplbond.CCH2NR2 (NR2 = morpholino, piperidino, NPh2, N(CH2Ph)2] were synthesized in 75-88% yield from formalin,

Me(CH2)16CO2CH2C.tplbond.CH, and the corresponding amine in dioxane by the

use of Cu(OAc)2. 29237-96-5P ΙT

CORPORATE SOURCE:

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 29237-96-5 CAPLUS

Octadecanoic acid, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX

L4 ANSWER 21 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1972:108230 CAPLUS

DOCUMENT NUMBER: 76:108230

ORIGINAL REFERENCE NO.: 76:17421a,17424a

4-Amino-2-buten-1-ol esters TITLE:

Willette, Robert E.; Driscoll, Richard C. AUTHOR (S): Sch. Pharm., Univ. Connecticut, Storrs, CT, USA CORPORATE SOURCE:

Journal of Medicinal Chemistry (1972), 15(1), 110-12 SOURCE:

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

Trans-4-amino-2-butene-1-ol esters, R2NCH2CH:CHCH2O2CR1, where R2N = Me2N or morpholino and R1 = Me or iso-Pr, were prepared by condensation of the desired amine with 4-chloro-2-butyn-1-ol, followed by LiAlH4 reduction

and esterification. The corresponding cis isomers were prepared by esterification of the aminobutynol followed by catalytic reduction with

Pd/C. None of these compds. (100mg/kg/week, 8 weeks) showed any hepatotoxicity in mice.

35956-47-9P 35956-48-0P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 35956-47-9 CAPLUS

2-Butyn-1-ol, 4-(4-morpholinyl)-, acetate (ester) (9CI) (CA INDEX NAME)

35956-48-0 CAPLUS

Propanoic acid, 2,2-dimethyl-, 4-(4-morpholinyl)-2-butynyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1969:447998 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 71:47998

ORIGINAL REFERENCE NO.: 71:8815a,8818a

Acetylene compounds of potential pharmacological TITLE: value. XII. Central and peripheral anticholinergic

activity of tertiaryaminoalkynyl esters of some

carboxylic acids

Dahlbom, Richard; Erbing, Birgitta; Olsson, Kerstin; AUTHOR(S):

George, Robert; Jenden, Donald J.

CORPORATE SOURCE: Farm. Fak., Stockholm, Swed. SOURCE:

Acta Pharmaceutica Suecica (1969), 6(3), 349-58 CODEN: APSXAS; ISSN: 0001-6675

DOCUMENT TYPE: Journal LANGUAGE: English

tert-Aminoalkynyl esters of 1-phenylcyclopentanecarboxylic acid, BΑ 1-phenyl-cyclohexanecarboxylic acid, and benzilic acid were more active than the esters of diphenylacetic acid and phenothiazine-10-carboxylic acid when tested for antagonist activity toward acetylcholine on isolated guinea pig ileum and for mydriatic activity in intact mice. Generally

esters of benzilic acid appeared to have the highest potency. The most effective of these compds. was about half as active as atropine in blocking the central effects of oxotremorine and its effect on contractions of the guinea pig ileum induced by acetylcholine was .apprx.14% that of atropine.

IT 24642-37-3 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmacology of)

24642-37-3 CAPLUS

Cyclopentanecarboxylic acid, 1-phenyl-, 4-morpholino-2-butynyl ester CN (8CI)

(CA INDEX NAME)

L4 ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1968:49456 CAPLUS DOCUMENT NUMBER: 68:49456 ORIGINAL REFERENCE NO.: 68:9563a,9566a TITLE: 4-Dialkylamino-2-butynyl-1phenylcycloalkanecarboxylates INVENTOR (S)': Dahlbom, Richard PATENT ASSIGNEE(S): Aktiebolag Astra SOURCE: U.S., 2 pp. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE PATENT NO. KIND DATE APPLICATION NO. ______ 19640716 US 3317526 19670502 US GI For diagram(s), see printed CA Issue. AB The title compds, exhibit tremorolytic action with a min. of side-effects, and thus are effective in therapy of Parkinson's disease. They are made preferably via the Mannich reaction, but alternative routes are possible. Thus, a mixture of 35 g. 1-phenyl-1-cyclopentanecarbonyl chloride (I) and 10 g. propargyl alc. was refluxed 15 min. and fractionated in vacuo to give propargyl 1-phenyl-1-cyclopentanecarboxylate (II), b0.3 107-8°. A solution of 10 g. II, 3.4 g. pyrrolidine, 1.6 g. (CH2O)n, and 0.15 g. CuCl in 30 cc. dioxane was refluxed 10 min., treated with 150 cc. H2O, extracted with Et20 (extract discarded), and alkalized with 5N NH4OH. The precipitated amino ester was taken up in Et20, the solution dried, and the HCl salt precipitated with HCl in Et20 to give 4-pyrrolidino-2 - butynyl 1 - phenyl - 1 cyclopentanecarboxylate (III).HCl, m. 105-7° (2:1 EtOH-Et20). Similarly prepared were: 4-diethylamino-2-butynyl 1-phenyl-1cyclopentanecarboxylate-HCl (IV), m. 93-4°; 4-diethylamino-2butynyl 1-phenyl-1-cyclohexanecarboxylate-HCl, m. 126-8°; and 4-piperidino-2-butynyl 1-phenyl-1-cyclopentanecarboxylate-HCl, m. 124-6°. A solution of 11.5 g. I, 7 g. 4-diethylamino-2-butyn-1-ol (V), and 6 g. NEt3 in 75 cc. C6H6 was refluxed 2 hrs., cooled, and filtered, the filtrate worked up, and the product treated with HCl in Et 20 to give IV. Similarly prepared were: 4-pyrrolidino-2-butynyl 1-phenyl-1-cyclohexanecarboxylate-HCl, m. 127-9°; 4-morpholino-2-butynyl 1-phenyl-1-cyclopentanecarboxylate-HCl, m. 123-5°. To 7 g. V were added 0.5 g. Na and 2.5 g. methyl 1-phenyl-1-cyclopentanecarboxylate (VI) and the mixture heated 3 hrs. at 50°/10 mm., thus removing MeOH as formed. After dilution with 100 cc. H2O and acidification to pH 5, unchanged VI was extracted with 50 cc. Et20, the solution alkalized and extracted with Et20, and IV isolated as above. 17781-98-5P RL: SPN (Synthetic preparation); PREP (Preparation)

L4 ANSWER 25 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1967:18418 CAPLUS DOCUMENT NUMBER: 66:18418 ORIGINAL REFERENCE NO.: 66:3523a,3526a Mannich reaction with propargyl alcohol TITLE: Salvador, Romano L.; Simon, D. AUTHOR (S): CORPORATE SOURCE: Univ. Montreal, Montreal, Can. Canadian Journal of Chemistry (1966), 44(21), 2570-5 CODEN: CJCHAG; ISSN: 0008-4042 Journal DOCUMENT TYPE: English LANGUAGE: CASREACT 66:18418 OTHER SOURCE(S): GI For diagram(s), see printed CA Issue. AB A group of aminobutynols (e.g. I. of the type R2NCH2C.tplbond.CCH2OH were prepared from propargyl alc. by the Mannich reaction, using CuSO4 as catalyst, and the probable course of reaction discussed. The effect of on the yield in the reaction was studied showing that the reaction should be run in a medium which is acidic enough to form and stabilize the postulated carbenium ion R2NCH2+ but not acidic enough to prevent the formation of Cu acetylide. 14597-23-0P 14597-33-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 14597-23-0 CAPLUS

Benzilic acid, 4-morpholino-2-butynyl ester (7CI, 8CI) (CA INDEX NAME)

RN 14597-33-2 CAPLUS
CN 1-Naphthalenecarbamic acid, 4-morpholino-2-butynyl ester (8CI) (CA INDEX NAME)

L4 ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)
(prepn. of)
RN 17781-98-5 CAPLUS
CN Cyclopentanecarboxylic acid, 1-phenyl-, 4-morpholino-2-butynyl ester hydrochloride (7CI, 8CI) (CA INDEX NAME)

● HCl

ORIGINAL REFERÊNCE NO.: 63:13201d-f TITLE: Anticholinergic agents-esters of 4-dialkyl (or 4-polymethylene)amino-2-butynols AUTHOR(S): Majewski, Robert F.; Campbell, Kenneth N.; Dykstra, Stanley; Covington, Robert; Simms, CORPORATE SOURCE: Mead Johnson Res. Center, Evansville, IN SOURCE: Journal of Medicinal Chemistry (1965), 8(5), 719-20 CODEN: JMCMAR; ISSN: 0022-2623 DOCUMENT TYPE: Journal LANGUAGE: English 4-Dialkyl (or 4-polymethylene) amino-2-butynols (I) were prepared from 4-chloro-2butynol and the corresponding secondary amines (Biel, at al., 52, 6335g). Four procedures were used for the preparation of the title esters RCO2CH2C.tplbond.CCH2R' (I): from propargyl alc. analogous to that of Jones (J., et al., CA 42, 8774e); by ester-alc. interchange from a Me ester and the appropriate 4-amino-2-butynol; by esterification of the aminobutynol with an acid chloride; and by ester-ester interchange from e.g. 4-diethylamino-2-butynyl acetate and the Me ester of an appropriate carboxylic acid. The compds. were tested for smooth muscle depressant, local anesthetic, and (or) anticholinergic actions. I.HCl (R = SMe, R' piperidino) was found to have local anesthetic activity equivalent to lidocaine hydrochloride. I.HCl [R = ZPhCOH (Z = cyclohexyl), R' = NEt2] was found to possess about 10% of the activity of atropine on several types of extravascular smooth muscle plus strong papavarine-like action. 3512-26-3P, 1-Naphthoic acid, 4-morpholino-2-butynyl ester, hydrochloride 3512-28-5P, 2-Naphthoic acid, 4-morpholino-2butynyl ester, hydrochloride 3512-36-5P, Acetic acid, (methylthio)diphenyl-, 4-morpholino-2-butynyl ester, hydrochloride RL: PREP (Preparation) (preparation of) 3512-26-3 CAPLUS 1-Naphthoic acid, 4-morpholino-2-butynyl ester, hydrochloride (7CI, 8CI) (CA INDEX NAME)

L4 ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

63:71627

ACCESSION NUMBER:

DOCUMENT NUMBER:

1965:471827 CAPLUS

● HCl

RN 3512-28-5 CAPLUS
CN 2-Naphthoic acid, 4-morpholino-2-butynyl eater, hydrochloride (7CI, 8CI)
(CA INDEX NAME)

ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

● HC1

RN 3512-36-5 CAPLUS Acetic acid, (methylthio)diphenyl-, 4-morpholino-2-butynyl ester, hydrochloride (7CI, 8CI) (CA INDEX NAME)

● HCl

L4 ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1964:16636 CAPLUS DOCUMENT NUMBER: 60:16636 ORIGINAL REFERENCE NO.: 60:2909d-h,2910a-c TITLE: Aminoacetylenes PATENT ASSIGNEE(S): Mead Johnson & Co. SOURCE: 7 pp. DOCUMENT TYPE Patent Unavailable LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. 19610725 GB 940540 GB 1961-26864 19631030 DE 1216866 DE 19610620 19650330 US 1961-118261 US 3176019 19600726 PRIORITY APPLN. INFO.: US

A solution of 1.56 g. paraformaldehyde (I) and 2.0 g. Me2NH in 10 ml. dry dioxane was allowed to stand at room temperature 10 min., 10 g. propargyl diphenylacetate in 25 ml. dry dioxane added, the mixture heated on a

BLEAM bath 17 hrs. under N and cooled slightly, the unreacted Me2NH removed, 2N HCl added, and the solution washed with Et20, cooled with crushed ice,

and made alkaline with 10% NaOH. The insol. oil was taken up in Et2O, the solution

dried (MgSO4) and filtered, dry HCl passed into the solution, and the filtered off to give 4-dimethyl-2-butynyl diphenylacetate-HCl, m.

180-1.5° (decomposition) (PrOH). Similarly prepared was 4-pyrrolidino-2-butynyl diphenylacetate-HCl, m. 140-2° (EtOAc-PrOH). Diphenylacetyl chloride (15 g.) was slowly added to 10.0

4-piperidino-2-butynol, b1.4 116°, n20D 1.5094 (prepared from 1-chloro-4-hydroxy-2-butyne and piperidine) dissolved in 30 ml. dry pyridine (exothermic reaction), the mixture heated on a steam bath 1 hr., cooled, and poured onto crushed ice-water, the mixture extracted with Et20, the

exts. washed with small portions 2N HCl to remove the residual pyridine, the Et20 solution washed with water and dried over MgSO4, and the product isolated by passing dry HCl into the filtered mixture to give 4-piperidino-2-butynyl diphenylacetate-HCl, m. 155-6.5° (EtOAc). To a solution of 17.2 g. a-chlorodiphenyl-acetyl chloride in 40 ml. dry pyridine was slowly added 7.0 g. 4-pyrrolidino-2-butynol (II), bl.0 98-104°, n20D 1.5055 (prepared from 1-chloro-4-hydroxy-2-butyne and pyrrolidine), and after the vigorous reaction subsided, the mixture

poured onto crushed ice-water, the aqueous solution extracted with Et2O, the exts. washed with water and extracted with 2N HCl, and the acidic exts. heated on a

steam bath 5 min., cooled, and made alkaline with 10% NaOH, the viscous oil taken up

in Et2O, the Et2O solution dried (MgSO4) and filtered, and the Et2O evaporated to give a yellow solid, which was triturated with Et20 to give

4-pyrrolidino-2-butnyl benzilate, m. 108-11.5° (aqueous EtOH) (HCl salt m. 132.5-4.5°)'. Diphenylisobutyryl chloride (18.1 g.) and 21.0 g.

L4 ANSWER 27 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1964:52495 CAPLUS DOCUMENT NUMBER: 60:52495 ORIGINAL REFERENCE NO.: 60:9191a-f Acetylene compounds of potential pharmacological TITLE: value. III. 4-Dialkylamino-2-butynyl esters of benzilic acid Dahlbom, Richard; Hansson, Birgitta; Mollberg, Rene AUTHOR (S): CORPORATE SOURCE: Kungl. Farm. Inst., Stockholm SOURCE: Acta Chemica Scandinavica (1963), 17(8), 2354-6 CODEN: ACHSE7; ISSN: 0904-213X DOCUMENT TYPE: Journal LANGUAGE: English OTHER SOURCE(S): CASREACT 60:52495 AB cf. CA 59, 8729h. The title compds. were prepared by the method of King Holmes (CA 41, 5121g). Low yields are obtained by trans esterification Me benzilate with the appropriate 4-dialkylamino-2-butyn-1-ol. The following Ph2CRCO2CH2C.tplbond.CCH2NR12 were prepared (R. R1, derivative, * yield, and m.p. given): Cl, Et, HCl salt, 81, 96-7°; HO, Et, HCl salt, 55, 127-8*, HO, Et, MeBr, 78, 149-50.5*; Cl, (NR12 =) pyrrolidino, HCl salt, 78, 164-5.5°; HO, (NR12 =) pyrrolidino, HCl salt, 86, 137-8° (base m. 110-12°); Cl. (NR12 *) piperidino, HCl salt, 76, 141-2°; HO, (NR12 =) piperidino, HCl salt, 52, 146-7°; HO, (NR12 +) morpholino, HCl salt, 50, 148-9°. The compds. had anticholinergic activity and inhibited tremors due to oxotremorine. 95130-66-8P, Benzilic acid, 4-morpholino-2-butynyl ester, hydrochloride RL: PREP (Preparation) (preparation of) 95130-66-8 CAPLUS

Benzilic acid, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA

INDEX NAME)

● HCl

ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN Et3N were cautiously mixed with 85 ml. anhyd C6H6, 10.1 g. II dissolved

20 ml. anhyd. C6H6 added dropwise, the mixt. heated on a steam bath 3 hrs., and the product isolated as in the previous example to give 4-piperidino-2-butynyl diphenylisobutyrate-HCl, m. 156.5-8.5° (C6H6). To a soln. of 9.5 g. Me u-methylthiodiphenylacetate and 4.9 g.II in 150 ml. n-heptane was added about 50 mg. NaOMe, the mixt. stirred and refluxed, the MeOH-n-heptane azeotrope collected to a total of 0.85 ml. (addnl. NaOMe added during the distn.), the mixt. cooled and poured onto crushed ice-water, the org. layer sepd., washed with water, and

with 2N HCl, and the acidic exts. washed with Et20 and made alk. with 10% NaOH. The free base was taken up in Et20, the soln, washed with water, dried (MgSO4), and filtered, and dry HCl passed in to give 5.4 g. 4-pyrrolidino-2-butynyl-α-methylthiodiphenylacetate-HCl, m. 154-6° (iso-PrOH). Similarly prepd. were (HCl salt m.p. given): 4-morpholino-2-butynyl benzilate, 158-60°; 4-diethylamino-2-butynyl- α -methylthiodiphenylacetate, 146-8° 4-piperidino-2-butynyl α-methylthiodiphenylacetate, 171.5-73°; 4-morpholino-2butynyl a-methylthiodiphenylacetate, 171-3.5°; 4-diethylamino-2-butynylphenyl-α-thienylglycolate (81.5-3.5*), 4-diethylamino-2-butynyl phenylcyclohexylglycolate, 129-30°; 4-dimethylamino-2-butynyl benzilate, 130-3°; 4-diethylamino-2-butynyl benzilate, 128.5-30.5*; 4-piperidino-2-butynyl benzilate, 141.5-4°; 4-piperidino-2-butynyl α-methoxydiphenylacetate, 170.5-72°; and 4-piperidino-2butynyl a-ethoxydiphenylacetate, 173.5-75°. A mixt. of 394.2 g. Me phenylcyclohexylglycolate and 293.1 g. 4-diethylamino-2-butynyl acetate was dissolved in 2.6 l. n-heptane by warming, the soln. heated with stirring to 60-70°, 8.0 g. NaOMe added, the temp, raised until the solvent distd., the distn. continued until no more MeOAc distd., the mixt. cooled to room temp., washed with water, and extd. with 165 ml. 2N HCl, the aq. exts. stirred to permit crystn. of the HCl salt, and crystn. completed by cooling to give 323 g. 4-diethylamino-2-butynyl phenylcyclohexylglycolate-HCl. A mixt. of 11.4 g. achlorodiphenylacetyl chloride and 4.9 g. 4-dimethylamino-2-butynol was heated 25 min. at 100-5°, then 30 min. at 70°, the oil washed thoroughly with anhyd. Et20 and dissolved in 100 ml. anhyd. EtOH, the soln. refluxed 25 hrs. with 5 g. Na2CO3, the mixt. cooled, filtered, and made basic with 10% NaOH, most of the EtOH removed in vacuo, the aq. mixt. extd. with Et20, the exts. washed with water and dried over MgSO4, and then anhyd. HCl passed into the Et20 soln. to give 4.0 g. 4-dimethyl-2-butynyl a-ethoxydiphenylacetate-HCl, m. 166.5-8.5°.

3512-36-5P, Acetic acid, (methylthio)diphenyl-, 4-morpholino-2-butynyl ester, hydrochloride 14597-23-0P, Benzilic acid, 4-morpholino-2-butynyl ester 95130-66-8P, Benzilic acid, 4-morpholino-2-butynyl ester, hydrochloride RL: PREP (Preparation) (preparation of)

3512-36-5 CAPLUS

Acetic acid, (methylthio)diphenyl-, 4-morpholino-2-butynyl ester,

hydrochloride (7CI, 8CI) (CA INDEX NAME)

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L4 ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

• HCl

RN 14597-23-0 CAPLUS CN Benzilic acid, 4-morpholino-2-butynyl ester (7CI, 8CI) (CA INDEX NAME)

RN 95130-66-8 CAPLUS
CN Benzilic acid, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA

• HCl

L4 ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

• HC1

RN 98075-12-8 CAPLUS
CN Carbanilic acid, 2,6-dimethyl-, 4-morpholino-2-butynyl ester,
hydrochloride (7CI) (CA INDEX NAME)

• HC1

RN 98222-92-5 CAPLUS
CN Carbanilic acid, 2-chloro-6-methyl-, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA INDEX NAME)

● HC1

RN 98249-62-8 CAPLUS
CN Carbanilic acid, o-methyl-, 4-morpholino-2-butynyl ester, hydrochloride
(7CI) (CA INDEX NAME)

L4 ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1963:448342 CAPLUS DOCUMENT NUMBER: 59:48342 ORIGINAL REFERENCE NO.: 59:8730b-d TITLE: Acetylene compounds of potential pharmacological value. II. 4-Amino-2-butynyl esters of phenylcarbamic acide. Dahlbom, Richard; Mollberg, Rene AUTHOR (S): CORPORATE SOURCE: Roy. Inst. Pharm., Stockholm SOURCE: Acta Chemica Scandinavica (1963), 17, 1182-3. CODEN: ACHSE7; ISSN: 0904-213X DOCUMENT TYPE: Journal LANGUAGE: English GI For diagram(s), see printed CA Issue. AB Et 2-chloro-6-methylphenylcarbamate (107 g.) was distilled in vacuo with (142 g.) to give 60 g. 2-chloro-6-methylphenylisocyanate, bl0 84-5°, n23D 1.5548. The appropriate phenylisocyanate (0.05 mole) and 0.05 mole IV were refluxed 3 hrs. in 25 ml. C6H6. The solution was cooled, diluted with Et2O, treated with ethereal HCl, and the precipitate was recrystd. (Et20-EtOH) and dried at 50°/0.05 mm. to give X. The following X were prepared (R1 R2, R, % yield, and m.p. given): H, H, V, 160.5-1.5°; Me, Me, V, 72, 182.5-3.5°; Me, Cl, V, 86, 171.2-2.5° (decomposition); H, H, VI, 54, 129.5-30.5°; Me, H, VI, B2, 129.5-31.0*; Me, Me, VI, 61, 175.5-6.5*; Me, Cl, VI, 86, 168-70° (decomposition); H, H, VII, 68, 148.5-9.5°; Me, H, VII, 79, 147.5-8° (decomposition); Me, Me, VII, 62,194-4.5° (decomposition); Me, Cl, VII, 77, 184.5-5* (decomposition); H, H, VIII, 174.5-5.5°; Me, H, VIII, 57, 177-8° (decomposition); Me, Me, VIII, 80, 192.5-3° (decomposition); Me, Cl, VIII, 77, 177-8° (decomposition); H. H. IX, 78, 152.5-3°; Ma, H. IX, 84, 173.5-4.5° (decomposition); Me, Me, IX, 73, 225-6° (decomposition); Me, Cl, IX, 75, 212-13° (decomposition). IT 97417-91-9P, 2-Butyn-1-ol, 4-morpholino-, carbanilate, hydrochloride 98075-12-8P, Carbanilic acid, 2,6-dimethyl-, 4-morpholino-2-butynyl ester, hydrochloride 98222-92-5P, Carbanilic acid, 2-chloro-6-methyl-, 4-morpholino-2-butynyl ester, hydrochloride 98249-62-8P, Carbanilic acid, o-methyl-, 4-morpholino-2-butynyl ester, hydrochloride RL: PREP (Preparation) (preparation of)

2-Butyn-1-ol, 4-morpholino-, carbanilate, hydrochloride (7CI) (CA INDEX

L4 ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

97417-91-9 CAPLUS

NAME)

● HC1

L4 ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 1963:448341 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 59:48341

ORIGINAL REFERENCE NO.: 59:8729h,8730a-b

Acetylene compounds of potential pharmacological TITLE: value. I, 4-Amino-2-butynyl esters of diphenylacetic acid, 1-phenylcyclopentane-1-carboxylic acid, and

phenothiazine-10-carboxylic acid AUTHOR (S): Dahlbom, Richard; Mollberg, Rene CORPORATE SOURCE: Roy. Inst. Pharm., Stockholm

Acta Chemica Scandinavica (1963), 17, 916-20 SOURCE:

CODEN: ACHSE7; ISSN: 0904-213X

DOCUMENT TYPE: Journal English LANGUAGE:

AB Esters of diphenylacetic acid (I), 1-phenylcyclopentane-1-carboxylic acid (II), and phenothiazine-10-carboxylic acid (III) with RCH2C.tplbond.CCH2OH

(IV) have been prepared, where R = NMe2 (V), NEt2 (VI), pyrrolidino (VII),

piperidino (VIII), and morpholino (IX). IV was obtained from C1CH2C.tplbond.CCH2OH and the appropriate amine by the method of Biel

et al., CA 52, 6335g). Reported were IV (R, % yield, b.p./mm., and n22D given): VII, 85, 112-13°/0.9, 1.5092; VIII, 71, 101-2°/0.4, 1.5043. A solution of 0.055 mole acid chloride, 0.05 mole IV, and 0.06

mole Et8N in 50 ml. C6H6 was refluxed 3-20 hrs., then cooled, filtered, and concentrated in vacuo. The residue was dissolved in 50 ml. Et20,

treated with HCl and the precipitate recrystd. from Et20-Et0H. Quaternary salts of III esters

were also prepared. The following RCH2C.tplbond.CCH2R1R2X were obtained (RH,

R1, R2X, % yield, and m.p. given): III, V, HCl, 48, 185-6° (decomposition); III, V, EtBr, 83, 158-9° (decomposition); III, VI, HCl.

181-2° (decomposition); III, VI, MeBr, 91, 141-2° (decomposition); III, VII, HCl, 69, 155.5-6.5° (decomposition); III, VII, MeBr, 89, 163-4° (decomposition); III, VIII, HCl, 72, 176-7° (decomposition); III, VIII, MeBr, 98, 170-1° (decomposition); III, IX, HCl, 64, 188-9° (decomposition); II, V, HCl, 86, 144-6°; II, VI, HCl, 57, 92.5- 4°; II, VIII, HCl, 65, 124-6°; II, IX, HCl, 71, 167-9°; I, VI, HCl, 79, 128-30°; I, VII, HCl, 83, 142-4°; I, VIII, HCl, 78, 158-60°; I, IX, HCl, 80, 160-1.5°.

97417-91-9 98075-12-8 98222-92-5 98249-62-8

(Derived from data in the 7th Collective Formula Index (1962-1966))

97417-91-9 CAPLUS

2-Butyn-1-ol, 4-morpholino-, carbanilate, hydrochloride (7CI) (CA INDEX NAME)

ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

● HCl

IT 17781-98-5P, Cyclopentanecarboxylic acid, 1-phenyl-, 4-morpholino-2-butynyl ester, hydrochloride 95130-43-1P, Acetic acid, diphenyl-, 4-morpholino-2-butynyl ester, hydrochloride 101318-97-2P, Phenothiazine-10-carboxylic acid, 4-morpholino-2-butynyl ester, hydrochloride RL: PREP (Preparation) (preparation of)

17781-98-5 CAPLUS Cyclopentanecarboxylic acid, 1-phenyl-, 4-morpholino-2-butynyl ester hydrochloride (7CI, 8CI) (CA INDEX NAME)

HCl

95130-43-1 CAPLUS Acetic acid, diphenyl-, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA INDEX NAME)

● HCl

(Continued) ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

● HCl

98075-12-8 CAPLUS Carbanilic acid, 2,6-dimethyl-, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA INDEX NAME)

98222-92-5 CAPLUS Carbanilic acid, 2-chloro-6-methyl-, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA INDEX NAME)

● HCl

Carbanilic acid, o-methyl-, 4-morpholino-2-butynyl ester, hydrochlorida (7CI) (CA INDEX NAME)

ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

101318-97-2 CAPLUS RN

Phenothiazine-10-carboxylic acid, 4-morpholino-2-butynyl ester, hydrochloride (7CI) (CA INDEX NAME)

HC1

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L4 ANSWER 31 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1961:24081 CAPLUS DOCUMENT NUMBER: 55:24081 55:4777£-g ORIGINAL REFERENCE NO.: Pharmacological studies on terpenes TITLE: Nishio, Hyoe AUTHOR (5): CORPORATE SOURCE: Med. Coll., Nara Nippon Yakurigaku Zasshi (1959), 55, 1552-67 SOURCE: CODEN: NYKZAU; ISSN: 0015-5691 DOCUMENT TYPE: Journal Unavailable LANGUAGE: AB The ganglionic blocking activities of terpenes containing quaternary radicals were studied. Introduction of isoketopinic acid group to 2-morpholinoethanol-MeI was effective not only in the potentiation but in the prolongation of the ganglionic blocking action of the morpholinium compound Ketopinic acid derivs. demonstrated but a transient blocking action. They were destroyed by human serums and guinea pig liver homogenates. On the other hand, isoketopinic acid derivs. and π -oxocamphor oxime deriva. were not destroyed, and showed a marked prolonged effect. 111357-35-8 (Derived from data in the 6th Collective Formula Index (1957-1961)) 111357-35-8 CAPLUS 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium iodide, 1,7-dimethyl-2-oxo-7norbornancarboxylate (6CI) (CA INDEX NAME)

ANSWER 32 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 2-carboxy-1,1-dimethylpiperidinium bromide (6CI) (CA INDEX NAME)

●a Br

ACCESSION NUMBER: 1961:22847 CAPLUS DOCUMENT NUMBER: 55:22847 ORIGINAL REFERENCE NO.: 55:4540a-c TITLE: Aminoalkynyl N-alkylpiperidinecarboxylates INVENTOR (S): Biel, John H. PATENT ASSIGNEE(S): Lakeside Laboratories, Inc. DOCUMENT TYPE: Patent LANGUAGE: Unavailable FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND PATENT NO. APPLICATION NO. DATE US 2867619 19590106 US 1956-620165 19561105 The title compds. RN. (CH2)4.CHCO2(CH2)nC.tplbond.C(CH2)nNR1R2 (I) when quaternized are useful as anti-hypertensive and ganglion-blocking agents. Morpholine (87 g.) in 135 cc. benzene was treated dropwise with a 41.8 g. 4-chloro-2-butyn-1-ol in 75 cc. benzene. After the exothermic reaction, the mixture was refluxed 3 hrs., cooled, filtered and distilled to give 90.8% 4-morpholino-2-butyn-1-ol (I), b0.01 104-6°. Me 2-(1-methylpiperidyl)carboxylate (31.4 g.), 31 g. I, and 0.5 g. NaOMe in 325 cc. heptane were heated and MeOH separated using a Dean-Stark tube to give 74.3% 4-morpholino-2-butynyl N-methylpipecolinate, b0.25 149-51* (short column); MeBr salt, m. 208-10°, yield 88.5%. Below are given other I prepared (R, NRIR2, & yield, b.p., n25D, and & yield and m.p. of the MeBr derivs. given): Me, NMe2, 70.2, b0.35 107-9*, 1.4824, 95.3, 193° (decomposition); Me, Et2N, 29.5, b0.5 133-5°, 1.4824, 60.4, 204-5° (decomposition); Me, pyrrolidino, 70.3, b0.55 138-9°, 1.4972, 98, 205° (decomposition); Me, morpholino, 74.3, b0.25 149-51°, 1.5012, 88.5, 208-10° (decomposition). 101261-21-6 (Derived from data in the 6th Collective Formula Index (1957-1961)) 101261-21-6 CAPLUS Pipecolic acid, 1-methyl-, 4-morpholino-2-butynyl ester (6CI) (CA INDEX

L4 ANSWER 32 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

L4 ANSWER 33 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1961:22846 CAPLUS DOCUMENT NUMBER: 55:22846 ORIGINAL REFERENCE NO.: 55:45391,4540a TITLE: 2-Chloropyridine 1-oxide INVENTOR(S): Shermer, David A. PATENT ASSIGNEE (S): Olin Mathieson Chemical Corp. DOCUMENT TYPE: Patent LANGUAGE: Unavailable FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

RL: PREP (Preparation) (preparation of) 109563-64-6 CAPLUS

109563-64-6P, 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with 2-carboxy-1,1-dimethylpiperidinium bromide

4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with

PATENT NO. APPLICATION NO. KIND DATE DATE 19600906 US 1958-772008 US 2951844 19581105 Aco2H (0.51 mole) as a 40% aqueous solution was added over 15 min. to 1 mole 2-chloropyridine (I) at 70°, the mixture stirred 150 min. at 70°, neutralized with NaOH, and the unreacted I distilled at about 115* with H2O. The distillate separated into 2 phases; 0.61 mole I was recovered by decantation, to leave a residue of 0.39 mole

2-chloropyridine 1-oxide (100 and 77% yields, based on I and AcO2H, resp.). The recovered I was recycled. 101261-21-6

(Derived from data in the 6th Collective Formula Index (1957-1961)) RN 101261-21-6 CAPLUS

Pipecolic acid, 1-methyl-, 4-morpholino-2-butynyl ester (6CI) (CA INDEX NAME)

IT

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L4 ANSWER 34 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                        1961:7981 CAPLUS
DOCUMENT NUMBER:
                         55:7981
ORIGINAL REFERENCE NO.: 55:1539h-i,1540a-d
                         Structure and reactions of gossypol
TITLE:
                         Shirley, David A.
AUTHOR (S):
CORPORATE SOURCE:
                         Univ. of Tennessee, Knoxville
                         Proc. Conf. Chem. Structure Reactions Gossypol
SOURCE:
                         Nongossypol Pigments Cottonseed, New Orleans (1959)
                         34-43
                         Journal
DOCUMENT TYPE:
                         Unavailable
LANGUAGE:
    For diagram(s), see printed CA Issue.
    Gossypol (I) anils were prepared from a series of aliphatic and aromatic
     amines to study the scope of the reaction and to prepare derivs. of I to
be
     used as (a) intermediates, (b) physiol. active compds., (c) dyes, or (d)
     model compds, for complexes of I with proteins. In group (a) were
     allylamine, diethylenetriamine, n-C18H37NH2, aminoacetal, p-H2NC6H4Ac,
     p-nitrobenzylamine, and p-BrC6H4CH2NH2, in group (b) H2NCH2CH2NMe2,
     p-H2NC6H4CO2Bu, p-H2NC6H4SO2NH2, and H2NCH2CH2Ph, in group (c)
     4-(o-tolylazo)-o-toluidine and p-H2NC6H4N:NPh, and in group (d)
     H2NCH2CO2Me, DL-lysine Me ester, and H2NCH2CONHCH2CO2Me. Deapogossypol
     hexa-Me ether was demethylated with C5H5N.HCl to deapogossypol, which was
     converted to the hexaacetate, deapogossypolone tetraacetate, and
     deapohydrogossypolone octaacetate. Anhydrogossypol was treated with
     cyclopentadiene to give a crystalline product of proposed structure II.
     Apogossypol was converted to the hexaallyl ether, which was heated in a
     mixture of Me2NPh and Ac2O to give the tetraacetate of a partly
     product. I was oxidized in 10% aqueous NaOH with 30% aqueous H2O2 at
     60-70° 8 min. to yield 2 crystalline compds., m. 231-3°, not
     further examined, and, m. 184-6°, tentatively identified by
     infrared and C-H analysis as
2,2'-dihydroxy-4,4'-diisobutyl-6,6'-dimethyl-
     biphenyl-3,3'-dicarboxylic acid (III); bis(2,4-dinitrophenyl hydrazide)
     269-70°. Ac20 with III gave a diacetate. A possible mechanism for
     the formation of III was given.
    109563-64-6
        (Derived from data in the 6th Collective Formula Index (1957-1961))
    109563-64-6 CAPLUS
    4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with
     2-carboxy-1,1-dimethylpiperidinium bromide (6CI) (CA INDEX NAME)
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ACCESSION NUMBER:
                         1961:7980 CAPLUS
DOCUMENT NUMBER:
                         55:7980
ORIGINAL REFERENCE NO. :
                        55:1539f-h
                         \beta-Aroylpropionic acids. XVII. Establishment of
TITLE:
                         the structure of \beta-(2-hydroxy-p-toluoy1)propionic
                         El-Abbady, A. M.; Baddar, F. G.; Labib, A.
AUTHOR (S):
CORPORATE SOURCE:
                         Ain-Shame Univ., Cairo
                         Journal of the Chemical Society (1960) 3420-1
SOURCE:
                         CODEN: JCSOA9; ISSN: 0368-1769
                         Journal
DOCUMENT TYPE:
                         Unavailable
LANGUAGE:
                         CASREACT 55:7980
OTHER SOURCE(S):
AB cf. CA 54, 22614i. The structure of \beta-(2-hydroxy-4-toluoy1)propionic
     acid (I), previously assumed on the basis of incomplete evidence (cf.
     Raval, et al., CA 33, 3779), was confirmed on the basis of the following
     reactions. I (5 q.) boiled 12 hrs. with 12 g. Me2SO4, 30 g. anhydrous
K2CQ3.
     and 15 ml. acetone gave 83% methyl \beta-(2-methoxy-4-toluoyl)propionate
     (II), m. 65-6° (C6H6-petr. ether). II (5.1 g.) boiled 2 hrs. with
     3% alc. KOH gave 4.4 g. \beta-(2-methoxy-4-toluoy1)propionic acid (III),
     m. 127-8° (C6H6). III (1 g.), 40 ml. 3% KOH, and 3 g. KMnO4 heated
     1 hr. on a boiling H2O-bath gave 0.6 g. 2-methoxyterephthalic acid. III
     (2 g.) reduced by the Martin modified Clemmensen method (30 hrs. at
     reflux) gave 1.8 g. γ-(2-methoxy-4-tolyl)butyric acid (IV), m.
     54-5° (petr. ether). IV (1 g.) refluxed 2 hrs. with 0.5 ml. POC13
     in 10 ml. tetrachloroethane, and the mixture hydrolyzed with cold H2O and
     then steam-distilled gave 5-methoxy-7-methyl-1-tetralone;
     2,4-dinitrophenylhydrazone, m. 223-4° (HOAc).
     109563-64-6
        (Derived from data in the 6th Collective Formula Index (1957-1961))
     109563-64-6 CAPLUS
     4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with
     2-carboxy-1,1-dimethylpiperidinium bromide (6CI) (CA INDEX NAME)
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●2 Br

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L4 ANSWER 36 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                         1960:34403 CAPLUS
DOCUMENT NUMBER:
                         54:34403
ORIGINAL REFERENCE NO.:
                         54:6789h-i,6790a-d
                         Camphor derivatives as the ganglionic blocking
TITLE:
agents.
                         I. Isoketopinic acid derivatives
AUTHOR (S):
                         Makaniani, Michio
CORPORATE SOURCE:
                         Yoshitomi Pharm. Inds., Ltd., Fukuoka-ken
SOURCE:
                         Yakugaku Zasshi (1959), 79, 1359-63
                         CODEN: YKKZAJ: ISSN: 0031-6903
DOCUMENT TYPE:
LANGUAGE:
                         Unavailable
    Na (1 mole) in liquid NH3 treated with 1 mole dialkylamino alc., the NH3
     replaced with 10 vols. PhMe, the solution treated with 1 mole
isoketopinoyl
     chloride, heated 1 hr. at 100°, cooled, the PhMe layer washed with
     NaHCO3, and the product distilled in vacuo gave dl-dialkylaminoalkyl
     isoketopinate (I). Me isoketopinate (1 mole) in 10 vols, heptane heated
     hrs. with 1.5 moles dialkylamino alc. and 0.1 mole MeONa, the solvent
     removed, the residue in C6H6 extracted with 5% HCl, the HCl layer
     and the oily product distilled gave I. I (1 mole) in C6H6 and 1.2 moles
     alkyl halide refluxed 5-7 hrs. and the product recrystd. (EtOH or Me2CO)
     gave I alkyl halide salt (II). I prepared were (dialkylaminoalkyl group,
     b.p./mm., m.p. of I.HCl, and m.p. of I alkyl halide salt given):
     Et2NCH2CH2, 160°/2, 145°, Et1, 152°; Me2N(CH2)3,
     171-3°/3, 162°, MeI, 181°; Me2NCH2CH2,
     142-6°/2, 191°, MeI, 208°; Et2N(CH2)3,
     155-7°/2, 167°, -, -; RCH2CH2(R = morpholino), 203°3,
     202°, MeI, 239° (BrCH2C.tplbond.CH malt m. 105°);
     R1CH2CH2 (R1 = piperidino), 170-6°/2, 247°, MeI,
     280°; R(CH2)3, 185-90°/2, 209°, MeI, 205°;
     R1(CH2)3, 180-5°/2, 203°, MeI, 190°; RCH2CHMe,
     150-7°/0.5, 183°, MeI, 208°; 4-
     ethoxyisoketopinoylpiperidinoethyl (III), 94°/0, 269°, MeI,
     193°; RCH2C.tplbond.CCH2, 170°/0.07, 165°, MeI,
     85°; R(CH2)4, 190-5°/0.9,103°, MeI, 65°.
     dl-Morpholinoethyl 3-chloroisoketopinate b0.6 172°; methiodide m.
     232°. Dialkylaminoalkylamine (1 mole), 1 mole isoketopinoyl
     chloride in 5 vols. CHCl3, and 1 mole C5H5N refluxed 3 hrs., the CHCl3
     layer extracted with 5% HCl, the HCl layer neutralized and the product
     recrystd. (C6H6) gave dl-N-isoketopinoyl-ω-dialkylaminoalkylamine
     (IV). IV (1 mole) in 10 vols. C6H6 heated with 1.2 moles MeI and the
     product recrystd. (EtOH) gave IV.MeI. IV prepared were
(dialkylaminoalkyl
    group, m.p., m.p. of IV.HCl, and m.p. of IV.MeI given):
     4-(2-hydroxyethyl)piperidino, 85°, 263°, 255°;
     Me2NCH2CH2, 48° 226°, 208°; Me2N(CH2)3, 85°,
     209°, 250°; Et2N(CH2)3, 59°, 174°,
     180°; R(CH2)3, 97°, 214°, 115°. I.MeI and
     hypotensive action (+ or ++) were (dialkylaminoalkyl group and activity
     given): Et2NCH2CH2, +; Me2N(CH2)3, +; Me2NCH2CH2, +; RCH2CH2, ++
     (CH.tplbond.CCH2Br salt, ++); R1CH2CH2, ++; R(CH2)3, ++.
     101865-08-1 111357-35-8
        (Derived from data in the 6th Collective Formula Index (1957-1961))
     7-Norbornanecarboxylic acid, 1,7-dimethyl-2-oxo-, 4-morpholino-2-butynyl
     ester, hydrochloride (6CI) (CA INDEX NAME)
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(Continued)

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● HCl

RN 111357-35-8 CAPLUS
CN 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium iodide,
1,7-dimethyl-2-oxo-7norbornancarboxylate (6CI) (CA INDEX NAME)

• I-

IT 101865-09-2P, 2-Butyn-1-ol, 4-morpholino-, 1,7-dimethyl-2-oxo-7-norbornanecarboxylate
RL: PREP (Preparation)
(preparation of)

RN 101865-09-2 CAPLUS
CN 7-Norbornanecarboxylic acid, 1,7-dimethyl-2-oxo-, 4-morpholino-2-butynyl ester (6CI) (CA INDEX NAME)

L4 ANSWER 37 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

• HCl

RN 111357-35-8 CAPLUS
CN 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium iodide,
1,7-dimethyl-2-oxo-7norbornancarboxylate (6CI) (CA INDEX NAME)

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L4 ANSWER 37 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                         1960:34402 CAPLUS
DOCUMENT NUMBER:
                         54:34402
ORIGINAL REFERENCE NO.: 54:6789e-h
TITLE:
                         Initiators and peroxide products of the liquid phase
                         autoxidation of 3-carene
                         Erofeev, B. V.; Chirko, A. I.
AUTHOR (S):
SOURCE:
                         Uchenye Zapiski, Belorus. Gosudarst Univ. im. V. I.
                         Lenina, Ser. Khim. (1956), 29, 15-22
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Unavailable
    The primary product of the autoxidn. of 3-carene (I) was I hydroperoxide
     (II). On reduction, II gave carenol. The other product of the
autoxidn. was
    2,2'-peroxide (III) of I. The following initiators were investigated:
     MnO, MnO2, Mn(HCO2)2 (IV), Mn(OAc)2 (V), Mn butyrate (VI), Mn atearate
     (VII), Fe203, Co304, (HCO2)2Co (VIII), Co(OAc)2 (IX), Co butyrate (X), Co
     stearate (XI), Co oxalate (XII), MoO3, WO3, PbO2, the hydrate of lead
     oxide (XIII), Pb(OAc)2 (XIV), SeO2, kaolin (XV), and montmorillonite
     (XVI). The best initiators were: MnO2, Fe2O3, Co3O4, MoO3, WO3, PbO2,
XV,
    XVI. Weak initiators were XII, XIII, and XIV. Se02 had an inhibiting
    effect. In the case of MnO, IV, VIII, XII, WO3, XIII, XIV, XV, or XVI,
    the amount of II found among the products of the autoxidn, approached
    autoxidn.; in the case of strong initiator, the amount of II was
     (35 g.) was oxidized in the presence of 1% XIV as long as the velocity of
     the oxidation began to decrease (4 1. O was necessary). The unchanged I
Was
    distilled and the residue fractionated 3 times in vacuo to give 8.5 g.
II,
    b0.024 49-50*, d20 1.0117, n20D 1.4991, MR 48.79, did not react
    with 2,4-dinitrophenylhydrazine, reacted violently with Pb(OAc)4 (XVII).
    The distillation residue of II contained III, b0.06 100°, d20 1.0717,
    n20D 1.5150, n50D 1.5050, gave no reaction with XVII. On reduction by
KI in
    AcOH II yielded carenol, b0.6 65-6°, d20 0.9892, n20D 1.4957,
    reacted with Na; phenylurethan m. 121°.
    101865-08-1 111357-35-8
        (Derived from data in the 6th Collective Formula Index (1957-1961))
    101865-08-1 CAPLUS
CN
    7-Norbornanecarboxylic acid, 1,7-dimethyl-2-oxo-, 4-morpholino-2-butynyl
     ester, hydrochloride (6CI) (CA INDEX NAME)
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L4 ANSWER 38 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
                         1958:35237 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         52:35237
ORIGINAL REFERENCE NO.:
                        52:6335g-i,6336a-d
TITLE:
                         Hypotensive agents. II. Aminoalkyl esters of
                         piperidinecarboxylic acids and their "reversed" ester
                         derivatives
AUTHOR (S):
                         Biel, John H.; Sprengeler, Edwin P.; Friedman, Harris
CORPORATE SOURCE:
                         Lakeside Labs., Inc., Milwaukee, WI
                         Journal of the American Chemical Society (1957), 79.
SOURCE:
                         6184-7
                         CODEN: JACSAT; ISSN: 0002-7863
                         Journal
DOCUMENT TYPE:
LANGUAGE:
                         Unavailable
OTHER SOURCE(S):
                         CASREACT 52:35237
   cf. C.A. 50, 2579d. Morpholine (87.0 g.) in 135 cc. C6H6 treated rapidly
    dropwise with 41.8 g. ClCH2C.tplbond.CCH2OH in 75 cc. C6H6, refluxed 3
    hrs., cooled, and filtered, the residue washed with C6H6, and the
combined
    filtrates distilled yielded 56.3 g. 4-morpholino-2-butyn-1-ol (I),
    oil, b0.1 104-6°, n25D 1.5087. Me N-methylpipecolinate (31.4 g.)
    and 31.0 g. I in 325 cc. heptone refluxed with 0.5 g. NaOMe under a
    Dean-Stark trap in which the liberated MeOH seps. from the heptane (2
    addnl. 0.3-g. portions NaOMe may be required to complete the reaction),
     50% of the heptane distilled, and the residue chilled, filtered, and
    gave 41.6 g. 4-morpholino-2-butynyl N-methylpipecolinate (II), b0.25
    149-51°, n25D 1.5012. II (14.0 g.) in 80 cc. iso-PrOH refluxed 3
    hrs. with 19.0 g. MeBr, cooled, and filtered yielded 20.8 g. II.2MeI, m.
    208-10° (decomposition) (hot EtOH). Similarly were prepared the
following
    aminoalkyl esters of 1-methylnipecotic acid (aminoalkyl group, b.p./mm.
    ester, and m.p. of dimethobromide given): Me2N(CH2)2, 103-5*/4.0,
    230-2°; Me2N(CH2)3, 106-9°/1.0, 238-9°;
    2-morpholinoethyl, 116-20°/1.0, 235-6°; 2-pyrrolidinoethyl,
    103-5°/1.0, 209-11°; o-ClC6H4CH2MeN(CH2)2,
    148-53*/0.15, - (dimethiodide, 178-9*); 2-diethylaminoethyl
    1-ethylnipecotinate, 104-6°/2.0, 221-2°.
    3-Dimethylamino-2-propyl 1-methylnipecotinate, 137-41*/17,
    255-7°; 2-dimethylaminoethyl isonipecotinate, 95-6°/1.0,
    276-7°. The following aminoalkyl esters of 1-methylpipecolinic
    acid (same data given): Me2N(CH2)2, 145-7°/23,245°;
    Me2N(CH2)3, 156-8°/23, - (dimethiodide, 224-6°);
    2-morpholinoethyl, 136-8°/1.2, 233-5°; o-
    ClC6H4CH2MeN(CH2)2, 100-5°/0.5, - (dimethiodide, 185°);
    3-morpholinopropyl, 140-2°/0.8, - (dimethiodide, 169-71°).
    2-Dimethylaminoethyl nicotinate, 154-6°/19, 218-19°. The
    following 1,x-MeC5H9N(CH2)mO2C(CH2)nNR2 (III) (ring position, m, n, NR2,
    b.p./mm. of ester, and m.p. of dimethobromide or in parentheses of
    dimethiodide given): 2, 1, 2, morpholino, 140-1°/0.8,
    188-9*; 3, 0, 2, pyrrolidino, 105-8*/0.3, (165-6*);
    3, 0, 2, morpholino, 126-9°/0.7, (182-3°); 3, 0, 1, NMe2,
    136-8°/16, 183-4°; 3, 0, 2, Me2N, 158-60°/37,
     (194-5°); 3, 1, 1, Me2N, 147-8°/22, 232-3°; 4, 0, 1,
    Me2N, 110-13°/8, 263-4°; 3, 0, 2, o-ClC6H4CH2MeN, -,
     (125°). 3-(N-o-Chlorobenzylpiperidyl)3-
    morpholinopropionate (IV), 178-88°/0.05, 88°;
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- L4 ANSWER 38 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 3-pyrrolidinopropionate analog of IV, 169-72°/0.05, 82-4°. The following analogs of II (b.p./mm., n25D of ester, and m.p. of dimethobromide given): Me2N, 107-9°/0.35, 1.4824, 193°; Et2N, 133-5°/0.50, 1.4824, 204-5°; pyrrolidino, 138-9°/0.55, 1.4972, 205°. The 1 lowering of blood pressure with 1.0 mg./kg. intravenously and 10 mg./kg. orally in the normotensive dog and the duration of the effect are tabulated for the various bisquaternary compds. Several of the compds. displayed potent and sustained hypotensive properties. The structural features necessary for optimum hypotensive activity are discussed. IT 101261-21-6P, 2-Butyn-1-ol, 4-morpholino-, 1-methylpipecolate 109563-64-6P, 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with 2-carboxy-1,1-dimethylpiperidinium bromide RL: PREP (Preparation) (preparation of) 101261-21-6 CAPLUS Pipecolic acid, 1-methyl-, 4-morpholino-2-butynyl ester (6CI) (CA INDEX
- Me C-O-CH₂-C=C-CH₂-N
- RN 109563-64-6 CAPLUS
 CN 4-(4-Hydroxy-2-butynyl)-4-methylmorpholinium bromide, ester with 2-carboxy-1,1-dimethylpiperidinium bromide (6CI) (CA INDEX NAME)
- Me Me O Me C-O-CH₂-C-CH₂-T N O